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,	PATENTSCOPE® • About Patents	Results of searching in PCT for: (melatonin AND zolpidem): 251 records Showing records 1 to 25 of 251:	[Search	n Summary]
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	o Content	Refine Search		
	o Glossary	(melatonin AND zolpidem)	4	
	o National Office Databases	Title	Pub. Date Int. Class	App. Num
	o Terms and Conditions	1. (WO 2009/042440) 1-METHYL NICOTINAMIDE AND DERIVATIVES FOR TREATMENT		PCT/ US2008/076
	Technology Focus	OF GASTRIC INJURY The present invention is directed to nicostramice certifatives, and their use in treating gastron.		
	PCT Resources	tie hegat neather in marke in industrian administration and use and accord and	.ttsin og otserene.	
	Priority Documents	2. (WO 2009/042092) 2-ARYL OR HETEROARYL INDOLE DERIVATIVES		PCT/ US2008/010
	Data Services	The present invention provides 2-anyl or heteroanyl indole derivatives which are ASIC channel	I modulators, pharmaceutical	compositions
	Statistics	compounds, and methods of using them as therapeutic agents		
,	• Patent Law	3. (WO 2009/039461) N-SUBSTITUTED PIPERIDINE DERIVATIVES AS SEROTONIN	26.03.2009 C07D 211/58	PCT/
	Life Sciences	RECEPTOR AGENTS		US2008/077
	Meetings	Disclosed herein are isolated forms of the compounds of Formula (I), (II), (II), (IV) and (V), or polymorph, or ester thereof. Also disclosed are methods of inhibiting an activity of a serotronin		
	• Contact	polyticidati, of ester trailed into discussed are hermone of inflaming at activity of a sectional memors of allevialing of realing various disease contamions and side effects.	receptor, control mesons a	dii delizanza
	Related Links	(WO 2009/034380) PIPERIDINE DERIVATIVES AS AGONISTS OF MUSCARINIC RECEPTORS	19.03.2009 C07D 211/74	POT/ GB2008/050
	International Patent	Compounds of Formula (i), or pharmaceutically acceptable salts thereof, wherein R <sp>2<th></th><th></th></sp>		
	Classification	well as saits and pinarmaceurical compositions treationing the compounds are prepared. They a	ure userui in inerapy, in paino	Clar in the me
	Natural Language IPC Search	5. (WO 2009/027697) NON-AQUEOUS PHARMACEUTICAL COMPOSITIONS	05.03.2009 A61K 9/00	PCT/

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6. (WO 2009/024823) OXADIAZOLE DERIVATIVES AS DGAT INHIBITORS

The present invention provides a composition for intranasat delivery of a drug comprising: (i) the drug; and (ii) a non-aqueous vehicle or glycol and at least one additional solvent selected from N-methylpyrrolidone, propylene carbonate, dimethyl sulfoxide and at least one i

ester. (b) from about 40 to 100 % by volume of Pr-metrypyrrolidone; or (c) from about 40 to 100% by volume of dimetrys sullidation (DA

26 02:2009 C07D 211/40 PCT/

GB2008/002



activities regarding patents and

Disclosed herein is at least one cyclopropyl amide derivative of formula (f), at least one charmaceutical composition comprising at least derivative disclosed herein, and at least one method of using at least one cyclopropyl amide derivative disclosed herein for treating at it receptor associated condition therewith.

(WO 2009/020642) PYRIDINE CARBOXAMIDE OREXIN RECEPTOR ANTAGONISTS

12 02 2009 A01N 37/16 PCT/ US2008/009

The present invention is directed to pyridyl carboxamide compounds which are antagonists of crexin receptors, and which are useful in of neurological and psychiatric disorders and diseases in which crexin receptors are involved. The invention is also directed to pharmac comprising these compounds and the use of these compounds and compositions in the prevention or treatment of such diseases in wh involven

8. (WO 2009/020569) TREATMENT OF PSYCHOSIS WITH A 5HT2A ANTAGONIST AND A 12.02.2009 C12N 5/06 U\$2008/009 METABOTROPIC GLUTAMATE RECEPTOR AGONIST OR POTENTIATOR

The present invention is directed to the use of a 5-HT2A aniagonist and an mGluR2/3 agonist, an mGluR2 agonist or an mGluR2 poter osychosis, including schizophrenia or bipolar disorder.

9. (WO 2009/018824) USE OF A COMPOSITION COMPRISING AT LEAST ONE BETA-

12.02.2009 A61K 31/165 PCT/ DK2008/000

A composition comprising specific beta-blockers such as bisoprolol and neblyolol for the treatment of insermia and/or another sleep dis should be given in such an amount that it causes a less than 40 % decrease in the amount of aMT6s in complete nocturnal urin. The or comprisation treatment comprising a specific peta-blocker in comprisation with another known drug e.g., metatoriin with similar effect for

(WO 2009/017716) PULSATILE GASTRIC RETENTIVE DOSAGE FORMS

BLOCKER FOR THE TREATMENT OF SLEEP DISORDERS

PCT/ 05.02.2009 A61K 9/00 US2008/009

Dosage forms for delayed and pulsed release of the apeutic agents into the stomach are described. The dosage forms are gastric refer achieve release of the therapeutic agent into the stomach and upper gastrointestinal tract subsequent to administration of the dosage for particular use in administration of acid-labile active agents such as proton pump inhibitors, and in treating gastric acid secretion such as disease (GEHD) and hociumal acid preakthrough (IVAB).

11. (WO 2009/017452) NEW CRYSTALLINE FORMS OF 2 -HYDROXY- 3- [5- (MORPHOLIN- 05.02.2009 C07D 413/14 PCT/ SE2008/050 4- YLMETHYL) PYRIDIN-2-YL] IH- INDOLE- 5 -CARBONITRILE CITRATE

The present invention relates to new crystalline forms of 2-trydroxy-3-[5-(morpholin-4-yimethyl)pyridin-2-yi]1-disH</i>-indole-5-carbonitr Form E, respectively, a process for their preparations, pharmaceutical formulations containing said compounds and to the use of said a and particularly to GISA3 related conditions and discreers.

12. (WO 2009/011775) AMIDOETHYL ALKYLAMINO OREXIN RECEPTOR ANTAGONISTS 22.01.2009 A01N 43/64 US2008/008

The present invention is directed to amidoethylamine compounds which are antagonists of orexin receptors, and which are useful in the neurological and psychiatric disorders and diseases in which crexin receptors are involved. The invention is also directed to pharmacet comprising these compounds and the use of these compounds and compositions in the prevention or treatment of such diseases in wh involved.

13. (WO 2009/009015) QUINAZOLINONE T-TYPE CALCIUM CHANNEL ANTAGONISTS

15.01.2009 A01N 43/54 PCT/ US2008/008

The present invention is directed to quinazolinone compounds which are antagonists of T-type calcium channels, and which are useful prevention of disorders and diseases in which T-type calcium channels are involved. The invention is also directed to pharmacourtical o these compounds and the use of these compounds and compositions in the prevention or treatment of such diseases in which T-type of 14. (WO 2009/006403) NON-PEPTIDE MACROCYCLIC HISTONE DEACETYLASE (HDAC)

Compounds of Formula (f) or (fi), and methods of making and using thereof, are described herein, wherein M represents a macrolide or group, optionally containing one or more heteroations, wherein the carbon atoms and/or heteroations are in a linear and/or cyclic arran group, A is a linking group connected to D. B is an alkyt, wherein M represents a macrolide subunit, R-ssb1
 inseq 1.555.2-(355.2-3

Indole derivatives that are useful for treating pain, inflammation and other conditions are described. Certain of the compounds are benz

INHIBITORS AND METHODS OF MAKING AND USING THEREOF

(WO 2008/157740) FAAH INHIBITORS

receptors are involved.

08.01.2009 A61K

31/7056

24 12 2008 C07D 209/30 PCT/

PCT/ US2008/068

US2008/067

cenzovi derivatives: The compounds are substituted at least at the 3 position of the indole. 16. (WO 2008/155573) CINNOLINE COMPOUNDS FOR USE IN THE TREATMENT OF 24 12:2008 A61K 31/502 PCT/ GB2008/050 SCHIZOPHRENIA This invention relates to the use of compounds having the structural formula (I) below; and their pharmaceutically acceptable salts, taut hydrolysable precursors, compositions in treating schizophrenia. 17. (WO 2008/155572) FUSED QUINOLINE DERIVATIVES USEFUL AS GABA 24.12.2008 C07D 471/14 PCT/ G82008/050 MODULATORS This invention relates to novel compounds having the structural formula (I) below, and their pharmaceutically acceptable saits, tautome precursors, compositions and methods of use thereof, wherein Rksp>1k/sp>, Rksp>2k/sp>, Rksp>3k/sp>, Rksp>4k/sp>, Rksp>5k/sp defined in the specification. These novel compounds provide a treatment or prophylaxis of anxiety disorders, schizophrenia, cognitive d disproses 18. (WO 2008/150364) CYCLOPROPYL PYRROLIDINE OREXIN RECEPTOR ANTAGONISTS 11.12.2008 A01N 43/82 U\$2008/006 The present invention is directed to cyclopropyl profine bis-amide compounds which are antagonists of orexin receptors, and which are prevention of neurological and psychiatric disorders and diseases in which previn receptors are involved. The invention is also directed compositions comprising these compounds and the use of these compounds and compositions in the prevention or treatment of such d receptors are involved. 19. (WO 2008/148798) CONTROLLED RELEASE PHARMACEUTICAL COMPOSITIONS FOR 11.12.2008 A61K 9/20 EP2008/056 PROLONGED EFFECT Layered pharmaceutical composition suitable for oral use in the treatment of diseases where absorption takes place over a large part of The composition comprising A) a solid inner layer comprising i) an active substance, and ii) one or more disintegrants/exploding agents agents or a mixture thereof, the solid inner layer being sandwiched between two outer layers B1) and B2), each outer layer comprising soluble and/or crystalline polymer or a mixture of substantially water soluble and/or crystalline polymers, the polymer being a polyolycol nomopolymer having a MW of at least about 100,000 dations, and b) a copolymer having a MW of at i... 20. (WO 2008/147518) PYRIDYL PIPERIDINE OREXIN RECEPTOR ANTAGONISTS 04.12.2008 C07D 401/12 PCT/ US2008/006 The present invention is, directed to pyridyl piperidine compounds of formula (I) which are antagonists of orexin receptors, and which a prevention of neurological and psychiatric disorders and diseases in which orexin receptors are involved. The invention is also directed

compositions comprising these compounds and the use of these compounds and compositions in the prevention or treatment of such d

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21. (WO 2008/147314) SPIROCYCLOPROPYL PIPERIDINE DERIVATIVES

04.12.2008 C07D 405/04 PCT/

SE2008/050

US2008/062

Disclosed herein is at least one piperidine derivative, at least one pharmaceutical composition comprising at least one piperidine derivat treating at least one histamine H3 receptor associated condition merewith

22. (WO 2008/143856) OXO BRIDGED DIAZEPAN OREXIN RECEPTOR ANTAGONISTS

27.11.2008 A01N 43/62 PCT/ US2008/006

The present invention is directed to oxo bridged diazepan compounds which are antagonists of crexin receptors, and which are useful of neurological and psychiatric disorders and diseases in which crexin receptors are involved. The invention is also directed to pharmac comprising these compounds and the use of these compounds and compositions in the prevention or treatment of such diseases in white providers.

23. (WO 2008/137923) DROXIDOPA AND PHARMACEUTICAL COMPOSITION THEREOF FOR THE TREATMENT OF MOOD DISORDERS, SLEEP DISORDERS, OR ATTENTION

13.11.2008 A61K 31/138 PCT/

FOR THE TREATMENT OF MOOD DISORDERS, SLEEP DISORDERS, OF DEFICIT DISORDERS

The present invention provides pharmaceutical compositions comprising droxidopa alone, or in combination with one or more turther as treatment of conditions, such as mood disorders, steep disorders, or attention deficit disorders, in certain embodiments, the composition

treatment of conditions, such as mood disorders, sleep disorders, or attention deficit disorders. In cartain embodiments, the composition the invention comprise droxidopa and a compound selected from the group consisting of DOPA decarboxylase inhibiting compounds, continuity of the invention compounds, nonepinephrine reuptake inhibiting compounds, nonepinephrine reuptake inhibiting compounds, nonepinephrine reuptake inhibiting compounds, represented inhibiting compounds, nonepinephrine reuptake inhibiting compounds, represented inhibiting compounds, represented inhibiting compounds, represented inhibiting compounds, represented inhibiting compounds, nonepinephrine reuptake inhibiting compounds, represented inhi

24. (WO 2008/136756) PYRROLOPYRIMIDIN-7-ONE DERIVATIVES AND THEIR USE AS PHARMACEUTICALS

13.11.2008 C07D 487/04 PCT/ SE2008/050

Compounds of formula (I) or pharmaceutically acceptable salts thereof wherein A1, R2, R3, R4, and R9 and are as defined in the specific pharmaceutical compositions including the compounds are prepared. They are useful in merapy, in particular in the management of pal

25. (WO 2008/130571) NUCLEAR RECEPTOR BINDING AGENTS

30.10.2008 C07C 327/00 PCT/ US2008/004

The present invention relates to a novel class of selective estrogen receptor modulators (SERMs). The SERM compounds are applicab and/or treatment of a variety of diseases and conditions including prevention and treatment of cancers such as prostate and breast can related preseases not trashes or vasomotor symptoms, neurological dispress; cardiovascular disease and obesity.

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melatonin: 21695 occurrences in 2204 records. zolpidem: 6704 occurrences in 1111 records. (melatonin AND zolpidem). 251 records. Search Time: 0.97 seconds.

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